APPENDIX Ia: MOLECULAR GENETIC CHARACTERIZATION DATA FOR TRANSGENIC PLANTS INTENDED FOR UNCONFINED RELEASE

INTRODUCTION

In July 1998, regulatory officials of the Canadian Food Inspection Agency (CFIA), Health Canada and the United States Department of Agriculture's Animal and Plant Health Inspection Service (USDA-APHIS) met to compare, and harmonize where possible, aspects of molecular genetic characterization that are part of their review processes for transgenic plants. Agreement on common requirements and acceptable analytical approaches for molecular genetic characterization will facilitate the submission of supporting data by developers seeking regulatory approval to incorporate such plants into agricultural production or commerce in both countries. This Appendix is one of the outcomes of this meeting. The Appendix summarizes and identifies similarities and differences in the critical elements of the molecular genetic characterization of transgenic plants considered during the review process by these participating agencies. Molecular genetic characterization is only part of the information considered during assessments of such plants conducted prior to commercialization.

The scope of this document is limited to consideration of the transformation process and vectors used during transformation; the genetic material that was potentially delivered to the recipient plant; the identification, inheritance, and expression of the genetic material in the transgenic plant, and the production of new proteins and RNAs encoded by the introduced genetic material. This document does <u>not</u> address specific types of techniques nor quality assurance practices (e.g., good laboratory practices) that are used to generate molecular genetic characterization data.

The agencies found very substantial areas of agreement in the types of molecular genetic characterization data they require to be submitted and considered. In addition to the specific data sets reviewed, the participants of both countries reaffirmed that reviews are still conducted on a case-by-case basis which allows for reviewing additional or fewer data sets, depending upon the individual case and the regulatory authority of the individual agencies. The use of the word "may" in this document is intended to reflect some of this flexibility in determining when data sets will be considered as an appropriate part of the entire application package. Therefore, consultations between regulatory agencies and individual applicants are considered to be an important part of the overall application process in making such determinations.

The critical elements of the molecular genetic characterization of transgenic plants described below apply to the review process of the participating agencies in both Canada and the United States, except where noted. The contents of this document will be reviewed and amended as necessary by these agencies. The current version (V. 2) reflects changes made following consideration of comments submitted by Argentina's National Advisory Committee on Agricultural Biotechnology (CONABIA) on September 21, 2000 and January 5, 2001. The glossary which follows has been included to provide definition to certain terms within the context of this document.

GLOSSARY

carrier DNA DNA used to expedite the preparation or the transformation of genetic material

into a plant but which is itself not part of the construct.

coding region A DNA sequence which when transcribed contributes to the production of the

mature RNA, which may or may not be translated to produce a protein. Coding regions can include complete or truncated open reading frames (excluding introns) which may be translated to produce a protein or peptide or which may be intentionally engineered to be untranslatable, for example as with

antisense constructs or ribozymes.

construct An engineered DNA fragment (eg. plasmid) which contains, but is not limited

to, the DNA sequences to be integrated into a target plant's genome.

database citations Publicly accessible sources of nucleotide or protein sequence information.

Four commonly used databases and their website addresses are:

GenBank: An annotated collection of all publicly available DNA sequences maintained by the National Institute of Health (NIH). http://www.ncbi.nlm.nih.gov/Web/Genbank/index.html

DNA Data Bank of Japan: The officially certified DNA bank of Japan, which collects DNA sequences from researchers. http://www.ddbj.nig.ac.jp/fromddbj-e.html **EMBL Nucleotide Sequence**: A database of DNA and RNA sequences collected from the scientific literature, patent applications, and directly submitted from researchers and sequencing groups. http://www.ebi.ac.uk/ebi_docs/embl_db/ebi/topembl.html

The **SWISS-PROT Protein Sequence Data Bank**: A database of protein sequences produced collaboratively by Amos Bairoch (University of Geneva) and the EBI. http://www.ebi.ac.uk/~flang/sp/sp.html

insert

That part of a construct (see above) which is integrated into the recipient plant's genome.

non-coding region

DNA sequences which lie outside of an open reading frame and which are not translated to become part of a protein. These might include DNA sequences that function either in the plant or other hosts to regulate or influence the expression or processing of gene products (e.g. scaffold attachment regions, promoters, leader sequences, enhancers, introns, and terminators) or to faciliate replication, transposition, recombination, or cleavage of DNA (e.g. origins of replication, transposable element repeat sequences, T-DNA borders, *lox* sequences, polylinkers, etc.). Others may be sequences of no known function (e.g. plasmid backbone sequences).

stability

The ability of the transgenic trait to be expressed in the transformed plant line and plant lines derived therefrom in a consistent, reliable, and predictable manner.

trait(s)

The phenotypic characteristic(s) conferred to the recipient plant by the transgenic insert.

vector

An automonously replicating DNA molecule into which foreign DNA is inserted and then propagated in a host cell.

MOLECULAR GENETIC CHARACTERIZATION OF TRANSGENIC PLANTS

1 THE TRANSFORMATION SYSTEM

- 1.1 Description of the transformation method
- 1.1.1 Describe and provide references for the transformation method, e.g. *Agrobacterium*mediated transformation or direct transformation by methods such as particle
 bombardment, electroporation, polyethylene glycol (PEG) transformation of protoplasts,
 etc.
- 1.1.2 For direct transformation methods, describe the nature and source of any carrier DNA used.
- 1.1.3 For *Agrobacterium*-mediated transformation, provide the strain designation of the *Agrobacterium* used during the transformation process, and indicate how the Ti plasmid based vector was disarmed, and whether *Agrobacterium* was cleared from the transformed tissue.
- 1.1.4 For transformation systems other than *Agrobacterium*, provide the following information:
- 1.1.4.1 Does the system utilize a pathogenic organism or nucleic acid sequences from a pathogen?
- 1.1.4.2 How were any pathogenesis-related sequences removed prior to transformation?
- 1.1.4.3 Did the transformation process involve the use of helper plasmids or a mixture of plasmids? If so, describe these in detail.
- 1.2 Description of the genetic material potentially delivered to the recipient plant material (the modification/constructs).

- 1.2.1 Provide a summary of all genetic components which comprise the vector including coding regions, and non-coding sequences of known function (for example see Table 1). For each genetic component provide a citation where these functional sequences were described, isolated, and characterized (publicly available database citations are acceptable) and indicate:
- 1.2.1.1 The portion of the full sequence that has been inserted, and its size.
- 1.2.1.2 The location, order, and orientation in the vector.
- 1.2.1.3 The function in the plant.
- 1.2.1.4 The source (scientific and common, or trade name, of the donor organism).
- 1.2.1.5 If the genetic component is responsible for disease or injury to plants or other organisms, or if it is a known toxicant, allergen, pathogenicity factor, or irritant.
- 1.2.1.6 If the donor organism is responsible for any disease or injury to plants or other organisms, produces toxicants, allergens or irritants or is related to organisms that do.
- 1.2.1.7 If there is a history of safe use of the source organism or components thereof.
- 1.2.2 If there has been a significant modification that affects the amino acid sequence of genes designed to be expressed in the plant, provide the citation. If the modified amino acid sequence has not been published, provide the complete sequence highlighting the modifications. Modifications that affect only a few amino acids of a known sequence can simply be stated without providing the complete sequence. Indicate whether the modifications are known or expected to result in changes in post-translational modifications or sites critical to the structure or function of the gene product.
- 1.2.3 Provide a detailed map of the vector (see Figure 1) with the location of sequences described above that is sufficient to be used in the analysis of data supporting the characterization of the DNA, including as appropriate the location of restriction sites and regions used as probes and/or primers used for PCR.

2 INHERITANCE AND STABILITY OF INTRODUCED TRAITS WHICH ARE FUNCTIONAL IN THE PLANT

- 2.1 For plants which are either male or female fertile or both, provide data that demonstrates the pattern and stability of inheritance and expression of the new transgene traits. If the new trait can not be directly measured by an assay, it may be necessary to examine the inheritance of the DNA insert directly, and expression of the RNA.
- 2.2 For plants which are either infertile or for which it is difficult to produce seed (such as vegetatively propagated male-sterile potatoes), provide data to demonstrate that the transgene trait is stably maintained and expressed during vegetative propagation over a number of cycles that is appropriate to the crop.

3 CHARACTERIZATION OF THE DNA INSERTED IN THE PLANT

- 3.1 For <u>all</u> coding regions, provide data that demonstrate if complete or partial copies are inserted into the plant's genome. Coding regions may include truncated sense constructs, sequences engineered to be nontranslatable, antisense constructs, and constructs containing ribozymes, regardless of whether or not the coding region is designed or expected to be expressed in the transgenic plant. For Canadian submissions, information may be required indicating the number of copies which have been inserted, including integration of partial copies; and for allopolyploid plants, information indicating into which parental genome insertion has occurred.
- 3.2 For noncoding regions associated with the expression of coding regions:
- 3.2.1 Data should demonstrate whether or not plant promoters are inserted intact with the coding regions whose expression they are designed to regulate. These data are relevant to consideration of points 4.1 and 4.2 below.
- 3.2.2 DNA analysis may be necessary for introns, leader sequences, terminators, and enhancers of plant-expressible cassettes. DNA analyses may be presented in the form of Southern analyses, DNA sequencing, PCR analyses, or other appropriate information.

- 3.2.3 DNA analysis may be necessary for promoters and other regulatory regions associated with bacteria-expressible cassettes.
- For noncoding regions which have no known <u>plant</u> function and are not associated with expression of coding regions:
- 3.3.1 DNA analysis may be required for some sequences of known function (e.g., *ori V* and *ori-322*, *bom*, T-DNA borders of *Agrobacterium*, and bacterial transposable elements).
- 3.3.2 DNA analysis is not required for any remaining sequences of the plasmid backbone when the plasmid is well characterized.

4 PROTEIN AND RNA CHARACTERIZATION AND EXPRESSION

- 4.1 For all <u>complete</u> coding regions inserted, provide data that demonstrates whether the protein is or is not produced as expected in the appropriate tissues consistent with the associated regulatory sequences driving its expression (e.g., if the gene is inducible, determine if the gene is expressed in the appropriate tissues under induction conditions). For virus resistant plants where the transgenes are derived from a viral genome, in addition to transgene protein analysis, determine transgene RNA levels in tissues consistent with the associated regulatory regions driving expression of the transgene. The following exceptions also apply:
- 4.1.1 If the protein concentration is below the limits of detection, mRNA data may be substituted.
- 4.1.2 Protein analysis for products of genes used only as selectable markers may be waived under certain circumstances, e.g. when there is at least one complete copy of a selectable marker gene present and the effective expression of the selectable marker gene is verified by the process used to select the transformed tissue.

- 4.1.3 For plants modified to express nontranslatable mRNA, truncated sense constructs, antisense constructs, or constructs containing ribozymes, since the function of these genetic constructs is to specifically alter the accumulation of a specific mRNA or protein present in the transgenic plant, provide data on the level of the <u>target</u> protein only (e.g. native tomato fruit polygalacturonase would be the target protein of antisense polygalacturonase to achieve altered fruit ripening). If the target protein levels are below levels of detection, determine target mRNA levels.
- 4.2 When analysis indicates that the insert contains a fragment (as opposed to the "complete") of a coding region designed to be expressed in a plant, determine whether a fusion protein or a read-through could be produced and in which tissues it may be located. Such fragments may be detected for example by analysis of DNA or RNA or expression data such as western blots.
- 4.3 Protein or RNA characterization may not be required for <u>fragments of genetic constructs</u> not expected to be <u>functional in the plant</u> (e.g., fragments of selectable marker genes driven by bacterial promoters.)